REMARKS

In the Office Action mailed January 22, 2009, claims 33-36 were rejected under 35

U.S.C. §112, second paragraph, for the reasons set forth in the Detailed Action section of the

Office Action.

By the foregoing proposed claim amendments to claims 33-36, Applicants have addressed

the deficiencies which gave rise to the rejection of these claims under 35 U.S.C. §112, second

paragraph, and Applicants now believe that this ground for rejecting these claims has been

rendered moot.

Turning now to the prior art based rejections, claims 1-6 and 22-32 have been rejected

under 35 U.S.C. §103(a) as being unpatentable over Tamura et al. and Smith et al.

Claims 1-11 and 22-32 have been rejected under 35 U.S.C. §103(a) as being unpatentable

over Tamura et al. and Smith et al. in view of Nguyen.

Claims 1-15 and 22-32 have been rejected under 35 U.S.C. §103(a) as being unpatentable

over Tamura et al. and Smith et al. in view of Nguyen and Hubbell et al. Claims 1-11 and 16-32

have been rejected under 35 U.S.C. §103(a) as being unpatentable over Tamura et al. and Smith

et al. in view of Nguyen and Kuhla et al.

For the following reasons, Applicants traverse these prior art based grounds for rejecting

the claims of the application under 35 U.S.C. §103(a).

Subject of the Invention

The present invention is directed to a compound based on hyaluronic acid (HA),

wherein alcohol groups of hyaluronic acid are esterified with rhein as such or in derived form, or

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a salt thereof as set forth in claim 1, and to related processes, pharmaceutical compositions,

medicinal products and devices, biomaterials, and methods according to the present invention.

In particular, the claimed compound according to the present invention presents the

following unexpected, advantageous properties:

- Avoiding the drawbacks associated with oral administration of rhein;

Long-term shelf stability as compared to HA alone (in particular, no hydrolytic

degradation), as is documented under Example 5, 1);

Increased syringeability as compared to HA alone, as documented under Example

5, 2);

- Increased protective effect against IL-1 induced MMP expression as compared to

either rhein or HA alone (synergetic effect between rhein and HA moities), as

documented under Example 7.

The patentability of the present invention is supported by experimental data as presented

in Examples 1 to 7.

Rejection of Claims 1-6 and 22-32

Concerning the Examiner's rejection of claims 1-6 and 22-32 on the ground of

obviousness under 35 U.S.C. 103 (a) over Tamura et al. (EP 1082963) and Smith et al. (Arthritis

& Rheum., 1999) please consider the following.

a. Tamura et al.-EP 1082963

Tamura et al. is directed to conjugates of hyaluronic acid (HA), a derivative thereof or a

salt thereof, and a therapeutic agent, for joint diseases. In particular, Tamura et al. aims at

conjugates of a therapeutic agent for joint diseases and hyaluronic acid, a derivative thereof or a

salt thereof, which would be capable to localize and retain hydroxamic acid in a joint cavity.

Amendment After OA Mailed January 22, 2009 Serial No. 10/590,625 Tamura et al. is further directed to a method for preparing such a conjugate, which

comprises binding a site of a therapeutic agent for joint diseases that does not affect the activity

of the agent (e.g., amino group, a carboxyl group, a hydroxyl group, a thiol group, or the like) to

a carboxyl group, a hydroxyl group, or a functional group at the reducing end of hyaluronic acid,

a derivative thereof or a salt thereof [0057], [0072], [0073]. Tamura et al. generally describes the

possibility of activating the above sites either on a therapeutic agent or on the hyaluronic acid, a

derivative thereof or a salt thereof, to form various types of bonds such as amide bond, ester

bond, thioester bond, ether bond, imino bond, sulphide bond, etc. [0073], to form the conjugate.

When Tamura et al. specifically disclose the activation of a carboxyl group, they broadly

and generally disclose the activation of this group either on a therapeutic agent for joint diseases

or on HA, an HA derivative, or a salt thereof, with the use of a dehydrative condensation agent

(such as carbodiimides, phosphoniums, uroniums, and the like) to form an amide bond, an ester

bond, or a thioether bond [0075]. Furthermore, Tamura et al. fail to guide the ordinarily skilled

person to choose "a site of a therapeutic agent for joint diseases that does not affect the activity

of the agent" to attach HA.

Tamura et al. generally states that the mode of administration of the conjugate disclosed

herein is not particularly limited or suitable for any administration route, including oral route

[0086].

In addition, Tamura et al. teaches in the direction of using a HA conjugate where HA is

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conjugated with a therapeutic agent. In Tamura et al.'s conjugate, HA plays the role of a vehicle

for the therapeutic agent with the aim of retaining the said therapeutic agent with the hydroxamic

acid in a joint cavity to prolong the drug action at this specific site.

Contrary to the Examiner's assertions on page 3 of the Office Action mailed January 22,

2009, Tamura et al. fails to specifically teach the attachment of a therapeutic agent to HA, a

hydroxyl group, through the activation of a carboxylic group in the therapeutic agent.

Furthermore, as acknowledged by the Examiner, Tamura et al. fails to guide the

ordinarily skilled person towards the use of a specific therapeutic agent for joint diseases, in

particular rhein for the preparation of a conjugate [0034].

At most, Tamura et al. discloses metalloproteinase inhibitors (in particular hydroxamic

acids, i.e., compounds chemically completely different from rhein) as a specific therapeutic agent

category for joint diseases for the preparation of a conjugate.

Tamura et al. does not address in any way shelf stability or syringeability of HA, or

anticipate a potential synergetic effect between the HA drug carrier and the conjugated drug.

Therefore, Applicants respectfully assert that *Tamura et al.* does not anticipate or render

obvious the claimed invention.

b. Smith et al.

Smith et al. is directed to a study on the effect of diacerhein on early stages of damage in

a canine model of osteoarthritis (OA), when administered by oral route. Oral diacerhein is said to

show a slowing down effect on the progression of OA at early stages of the damage. However,

Smith et al. disclose that diacerhein does not have a significant effect at later stages of OA.

Smith et al. further disclose the absence of reduction in collagenase activity

(metalloproteinase) in OA cartilage following diacerhein treatment (p. 553, col. 2).

Smith et al. disclose that during the four-week treatment dogs suffered loose stools, a side

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effect of oral administration of diacerhein and rhein also mentioned in the present application

(see p. 1, [0005], as originally filed).

Smith et al. fail to teach any other administration route than oral route for diacerhein.

Smith et al. fail to teach any combination of diacerhein with any another agent, such as

HA.

Therefore, Applicants respectfully assert that Smith et al. does not anticipate or render

obvious the claimed invention.

Furthermore, when reading Tamura et al. in combination with Smith et al., the ordinarily

skilled person would have no guidance at all nor incentive to select any particular site of

conjugation on a therapeutic agent for joint diseases, with a specific site of HA, to form a

conjugate.

In particular, the ordinarily skilled person would not have any incentive to particularly

select rhein or a derived form, or a salt thereof, as a therapeutic agent for joint diseases to form a

conjugate for joint disease treatment.

Furthermore, Smith et al. fail to palliate the deficiencies of the teachings of Tamura et al.

To the contrary, when reading Smith et al., the person ordinarily skilled in the art would be

taught that diacerhein lacks inhibitory effect on collagenase activity (metalloproteinase) in OA,

and lacks significant effect at later stages of OA.

Tamura et al. and Smith et al., taken separately or in combination, do not provide any

teaching whatsoever of the present invention and its advantageous properties. Therefore,

Applicants respectfully assert that claims 1-6 and 22-32 are not anticipated by or rendered

obvious by Tamura et al. and Smith et al. taken separately or in combination.

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Rejection of Claims 1-11 and 22-32

Concerning the Examiner's rejection of claims 1-11 and 22-32 on the ground of

obviousness under 35 U.S.C. 103 (a) over Tamura et al. (EP 1082963) and Smith et al. (Arthritis

& Rheum., 1999) in view of Nguyen et al. (US 5,612, 321) please consider the following.

a. Tamura et al.-EP 1082963

Tamura et al. fail to render claims 1-6 and 22-32 obvious as discussed above. Further,

Tamura et al. fail to teach a process according to claims 7-11.

Therefore, Applicants respectfully assert that Tamura et al. does not render the claimed

invention obvious.

b. Smith et al.

Smith et al. fail to render obvious claims 1-6 and 22-32 as discussed above. Further,

Smith et al. fail to teach a process according to claims 7-11.

Therefore, Applicants respectfully assert that Smith et al. does not render the claimed

invention obvious.

c. Nguyen et al. –US 5,612, 321

Nguyen et al. relate to polysaccharides (including HA or crosslinked HA) grafted to

antioxidants on at least one hydroxyl group of the polysaccharide. Nguyen et al. aim at

generating polysaccharides with increased resistance to hydroxyl radicals by grafting them with

antioxidants.

Nguyen et al. do not teach a compound according to present claim 1. Further, Nguyen et

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al. do not teach the preparation of a conjugate of rhein or a derived form, or a salt thereof, nor

the preparation of a conjugate through a process according to present claim 7.

The process disclosed in Nguyen et al., which the Examiner cited (Example 6), relates to

the grafting of HA to an antioxidant by reacting an ammonium salt of HA with 3,5-di-t-butyl-4-

hydroxybenzoyl chloride in the presence of NMP (polar solvent) and ammonium ion,

exchanging the resulting product to sodium salt in aqueous solution.

Further, when reading Nguyen et al., the ordinarily skilled person would have no

incentive to replace the antioxidants by rhein for conjugation with HA.

Therefore, contrary to the Examiner's assertions on page 5 of the Office Action, Nguyen

et al. fail to specifically teach a process according to claim 7.

Therefore, the Applicants respectfully assert that Nguyen et al. does not render the

claimed invention obvious.

Furthermore, Nguyen et al. fail to palliate the deficiencies of the teachings of Tamura et

al. and Smith et al.

Therefore, based on the above reasons, Applicants respectfully assert that claims 1-11

and 22-32 are not rendered obvious by Tamura et al., Smith et al. or Nguyen et al. taken

separately or in combination.

Claims 1-15 and 22-32

Concerning the Examiner's rejection of claims 1-15 and 22-32 on the ground of

obviousness under 35 U.S.C. 103 (a) over Tamura et al. (EP 1082963) and Smith et al. (Arthritis

& Rheum., 1999) in view of Nguyen et al. (US 5,612, 321) and Hubbell et al. (US 5,834,274),

please consider the following.

a. Tamura et al.-EP 1082963

Tamura et al. fail to render obvious claims 1-15 and 22-32 as discussed above.

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Therefore, Applicants respectfully assert that Tamura et al. do not render the claimed

invention obvious.

b. Smith et al.

Smith et al. fail to render obvious claims 1-15 and 22-32 as discussed above.

Therefore, the Applicants respectfully assert that Smith et al. reference does not render

the claimed invention obvious.

c. Nguyen et al. –US 5,612, 321

Nguyen et al. fail to render obvious claims 1-11 and 22-32 as discussed above. Further,

as acknowledged by the Examiner, Nguyen et al. fail to teach the preparation of a HA conjugate

in a non polar aprotic solvent in the presence of a hydrogen ion acceptor according to the present

invention.

d. Hubbell et al. - US 5,834,274

Hubbell et al. relate to methods of coating and/or encapsulating surfaces and three-

dimensional objects with cross-linked networks of water soluble polymers. Hubbell et al.

discloses methods for polymerization of macromers using visible or long wavelength ultraviolet

light to encapsulate or coat either directly or indirectly living cells or tissues with polymeric

coatings.

First, Hubbel et al. deals with encapsulation and coating technologies that are far remote

from the field of the present invention.

Further, *Hubbell et al.* fail to teach the preparation of a conjugate of HA.

Hubbell et al. do not teach a conjugate of rhein or a derived form, or a salt thereof, nor

the preparation of a conjugate of the invention through a process according the invention, either.

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The Examples 1 and 2 from Hubbel et al. cited by the Examiner are directed to the

synthesis of PEG acrylates (diacrylate and tetraacrylate), which are far remote from the present

invention.

Furthermore, Hubbel et al. fail to palliate the deficiencies of the teaching of Tamura et

al., Smith et al. and Nguyen et al.

Therefore, based on the above reasons, Applicants respectfully assert that claims 1-11

and 22-32 are not rendered obvious by Tamura et al., Smith et al., Nguyen et al. and Hubbel et

al. taken separately or in combination.

Claims 1-11 and 16-32

Concerning the Examiner's rejection of claims 1-11 and 16-32 on the ground of

obviousness under 35 U.S.C. 103 (a) over Tamura et al. (EP 1082963) and Smith et al. (Arthritis

& Rheum., 1999) in view of Nguyen et al. (US 5,612, 321) and Kuhla et al. (US 4,788,187)

please consider the following.

a. Tamura et al.-EP 1082963

Tamura et al. fail to render obvious claims 1-15 and 22-32 as discussed above. Further,

Tamura et al. fail to teach acid chloride of rhein and a process for preparation thereof.

Therefore, Applicants respectfully assert that Tamura et al. reference does not render the

claimed invention obvious.

b. Smith et al.

Smith et al. fail to render obvious claims 1-15 and 22-32 as discussed above. Further,

Smith et al. fail to render obvious an acid chloride of rhein and a process for preparation thereof.

Therefore, Applicants respectfully assert that Smith et al. does not render the claimed

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invention obvious.

c. Nguyen et al. -US 5,612, 321

Nguyen et al. fails to render obvious claims 1-15 and 22-32 as discussed above.

Further, Nguyen et al. fail to teach the preparation of an acid chloride of rhein. In

particular, Nguyen et al. fails to teach the preparation of an acid chloride of rhein comprising

preparing a suspension of rhein in an aprotic non-polar solvent, and adding an amount of SOC12

so as to obtain a molar ratio between SOC12 and rhein greater than 10.

Therefore, the Applicants respectfully assert that Nguyen et al. does not render the

claimed invention obvious.

d. Kuhla et al. - US 4,788,187

Kuhla et al. relate to benzocyclobutene aminoalkylene ether and thioether compounds

and their use in gastrointestinal disorders.

Kuhla et al. do not render obvious a conjugate of rhein or a derived form or a salt thereof

nor the preparation of a conjugate of through a process according the present invention, either.

The Example 2 from Kuhla et al. cited by the Examiner is directed to the synthesis of a

benzocyclobutene derivative (5-(3-aminopropoxy)-1-(1-piperidinylmethyl)benzocyclobutene),

which is far remote from the present invention.

Therefore, Applicants respectfully assert that Kuhla et al. does not render the claimed

invention obvious.

Furthermore, Kuhla et al. fail to palliate the deficiencies of the teaching of Tamura et al.,

Smith et al. and Nguyen et al.

Therefore, based on the above reasons, Applicants respectfully assert that claims 1-11

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and 16-32 are not rendered obvious by Tamura et al., Smith et al., Nguyen et al. and Hubbel et

al. taken separately or in combination.

For all these foregoing reasons, Applicants respectfully request entry of the foregoing claim amendments, reconsideration of the present Application in light thereof, and in light of the foregoing remarks, and then allowance of all claims 1-36, as amended, over all the prior art of record.

Respectfully submitted,

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